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Dirhodium-Catalyzed Enantioselective B–H Bond Insertion of *gem*-Diaryl Carbenes: Efficient Access to *gem*-Diarylmethine Boranes

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Dedicated to 100th Anniversary of Chemistry at Nankai University

Abstract: The scarcity of reliable methods for synthesizing chiral gem-diarylmethine borons limits their applications. Herein, we report a method for highly enantioselective dirhodium-catalyzed B-H bond insertion reactions with diaryl diazomethanes as carbene precursors. These reactions afforded chiral gem-diarylmethine borane compounds in high yield (up to 99% yield), high activity (turnover numbers up to 14300), high enantioselectivity (up to 99% ee) and showed unprecedented broad functional group tolerance. The borane compounds synthesized by this method could be efficiently transformed into diaryl methanol, diaryl methyl amine, and triaryl methane derivatives with good stereospecificity. Mechanistic studies suggested that the borane adduct coordinated to the rhodium catalyst and thus interfered with decomposition of the diazomethane, and that insertion of a rhodium carbene (generated from the diaryl diazomethane) into the B-H bond was most likely the rate-determining step.

Introduction

Organoboron compounds are widely used in organic synthesis, materials science, medicine, and other fields.^[1-9] In particular, chiral gem-diarylmethine boron compounds, which have a unique gem-diaryl framework, are powerful synthons for the construction of bioactive compounds with a diaryl methane^[10-12] or triaryl methane^[12-16] motif via C-B bond transformations. Therefore, enantioselective synthesis of gem-diarylmethine boron compounds has attracted widespread interest, but it nevertheless remains a challenge. Since 2008, Aggarwal and co-workers^[17-19] have published several reports on lithiation-borylation reactions of chiral benzyl carbamates with aryl boronates for asymmetric synthesis of gem-diarylmethine borates containing a chiral quaternary carbon center (Scheme 1 A). Crudden and co-workers^[20] modified Aggarwal's protocol by adding an equivalent of chiral bisoxazoline ligands to prepare gem-diarylmethine boronic esters with a chiral tertiary carbon center, and these

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investigators used their protocol to synthesize chiral triaryl methane derivatives (Scheme 1 A). Recently, the groups of Liao^[21] and Tortosa^[22] independently developed coppercatalyzed 1,6-boration reactions of *para*-quinone methides for construction of chiral *gem*-diarylmethine borates (Scheme 1 B). To the best of our knowledge, this is the only catalytic asymmetric method for accessing *gem*-diarylmethine boron compounds. The use of strong bases in all the abovementioned methods strongly limits the functional group diversity of the *gem*-diarylmethine boron products. For instance, commonly encountered functionality such as esters, nitro groups, nitriles, and amides are incompatible with these methods. Moreover, for 1,6-boration of *para*-quinone me-

(A) Lithiation-borylation (Stoichiometric asymmetric synthesis)



(B) Cu-catalyzed 1,6-boration of para-quinone methides



(C) Rh-catalyzed B-H bond insertion of gem-diarylcarbene (This work)



Tolerated functional groups: F, Cl, Br, CF₃, OMe,

NO₂, CN, CO₂Me, CONMe₂, acetal, SO₂Me, etc.



OF

TON up to 14300 up to 99% yield up to 99% ee

Tolerated heteroaromatics: indole, benzofuran, pyridine *Scheme 1.* Methods for synthesis of chiral *gem*-diarylmethine boron compounds. thides (Scheme 1B), the steric requirements for the substrates substantially limit the structural diversity of the products.

We reasoned that B-H bond insertion reactions of carbenes might be useful for the synthesis of gem-diary-Imethine boranes. Transition-metal-catalyzed asymmetric B-H bond insertion reactions of carbenes have provided a new method for the synthesis of chiral organoboron compounds.^[23-33] Since we reported the first example of a copper-catalyzed asymmetric B-H bond insertion reaction with α -diazophenylacetate as a carbene precursor,^[25] catalytic asymmetric B-H bond insertion reactions have been successfully used for enantioselective construction of B-C bonds. Various carbene precursors such as a-diazophenylacetates,^[25,27,33] α -diazophenylketones,^[26,27] α -diazopropionates,^[30,32] trifluorodiazoalkanes,^[28] ene-yne-carbonyls,^[29] and tosylhydrazones^[31] have been used in these insertion reactions. Although with these progresses, the catalytic enantioselective B-H bond insertion is still in its infancy compared with the closely related well-established catalytic enantioselective C-H bond insertion.^[34-37] Thus, the use of B-H bond insertion in the synthesis of inconveniently available chiral organoboron compounds is highly desired. Herein, we report an efficient method for the synthesis of chiral gemdiarylmethine boron compounds by means of B-H bond insertion reactions of diaryl diazomethane compounds with catalysis by commercially available chiral dirhodium complexes (Scheme 1 C). Compared with the known methods for synthesis of gem-diarylmethine boron compounds, our method showed greatly enhanced functional group tolerance and thus enabled highly enantioselective synthesis of chiral gemdiarylmethine boron compounds with unprecedented structural diversity. Moreover, the compounds prepared in this study were bench-stable and could undergo several important functional group transformations with high preservation of stereochemistry, exhibiting great application prospects.

Results and Discussion

The highly enantioselective rhodium-catalyzed Si-H bond insertion reactions^[38,39] and cyclopropanation reactions^[40] using diaryl carbenes have been well established recently. Inspired by these pioneer works, we began our studies by carrying out reactions of 4-nitrophenyl phenyl diazomethane (1a) with trimethylamine-borane adduct 2a in the presence of 1 mol% of various chiral dirhodium catalysts in DCM at room temperature (Table 1). The B-H bond insertions were complete in minutes, giving desired gemdiarylmethine borane 3aa in moderate to high yields (entries 1–9). Of the tested catalysts, $Rh_2(S-TBPTTL)_4$ gave the highest yield and enantioselectivity (entry 4). Solvent screening revealed that other chlorinated solvents gave similar outcomes (compare entries 4, 10, and 11); whereas a coordinative solvent (THF) markedly decreased the reaction rate (entry 12), and a nonpolar solvent (toluene) slightly improved the enantioselectivity and the yield (entry 13). Lowering the reaction temperature improved the enantioselectivity further without compromising the yield (entries 13-16): 3aa was obtained in 96% yield and 91% ee at -40°C in toluene **Table 1:** Rh-catalyzed enantioselective B–H bond insertion of 4-nitrophenylphenyl diazomethane **1 a** with trimethylamine-borane adduct **2 a**: optimization of reaction conditions.^[a]



[a] Reaction conditions (procedure A): [Rh]/1a/2a = 0.002:0.2:0.24 (mmol), 2 mL solution of 1 a was dropped into a 2 mL solution of 2 a and [Rh]. All the reactions were completed within 3 min unless otherwise noted. Isolated yields were given. The *ee* values were determined by chiral HPLC, using chiral-phase IC-3 column. [b] Reaction time: 10 min. [c] Performed at 0°C. [d] Performed at -20°C. [e] Performed at -40°C.

(entry 16). We also tested numbers of chiral Cu^I and Rh^I catalysts in the template reaction but got unsatisfactory results (see Table S1 and Table S2 for details).

A series of borane adducts **2** were then evaluated in reactions with diazomethane **1a** (Table 2). Tertiary amineand phosphine-borane adducts smoothly underwent B–H bond insertion reactions and afforded the desired products in high yields with *ee* values similar to those obtained with **2a** (entries 1–5, 8, 9), whereas a borane adduct stabilized by a secondary amine failed to give any of the desired product due to catalyst decomposition (entry 6). 3,5-Dimethylpyridine-borane **2f** and N-heterocyclic carbene-borane **2i** also afforded high yields with moderate enantioselectivities under the standard reaction conditions (entries 7 and 10).

Next, we investigated $Rh_2(S$ -TBPTTL)₄-catalyzed asymmetric B–H bond insertion reactions of various diaryl diazomethanes 1 with amine-borane adduct **2a** (Scheme 2). The

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Research Articles

able 2: Rh-catalyzed asymmetric B—H bond insertion of 4-nitroph	e-
ylphenyl diazomethane 1 a with various borane adducts. ^[a]	

O ₂ N	N ₂ + 1a	H₃B - LB 2	Rh ₂ (S-TBPTTL), PhMe, -40 °C, 0	4 (1 mol %) 0.5 - 12 h O ₂ N	H ₂ B ^{LB}
Entry	LB		Product	Yield [%]	ee [%]
1	NMe ₃		3 aa	96	91
2	NEt₃ Me		3 ab	99	90
3			3 ac	95	92
4	Me		3 ad	97	91
5			3 ae	80	90
6	NHMe ₂	S	NA	NA	NA
7	Me	Me	3 af	95	76
8	P ⁿ Bu₃		3 ag	82	92
9	PMe ₂ Ph		3 ah	92	86
10	Me~N	N ^{-Me} /	3 ai	84	71

[a] Reaction conditions: $Rh_2(S-TBPTTL)_4/1a/2 = 0.002/0.2/0.24$ (mmol), 2 mL solution of 1a was dropped into a 2 mL solution of 2a and $Rh_2(S-TBPTTL)_4$. Isolated yields were given. The *ee* values were determined by chiral HPLC. NA = Not available.

reactions of all the tested diazo substrates smoothly gave the desired gem-diarylmethine borane compounds in good to excellent yields with high enantioselectivities (71-99% ee). Various functional groups, including nitro groups (3aa–3la), halogen atoms (3 fa-3 ha), trifluoromethyl groups (3 ia, 3 ta), an acetal (3la), an amide (3ma), an ester (3na), a sulfone (30a), a trifluoromethoxy group (3pa), and a cyano group (3 ga), were well tolerated under the reaction conditions. The enantioselectivity appeared to depend on electronic differences between the Ar¹ and Ar² groups of diaryl diazomethane substrates 1, where the electron donor and acceptor groups on each aryl ring play an important role. This phenomenon was consistent with previous studies.^[38-40] When Ar¹ had an electron-withdrawing para-nitro group, the enantioselectivity decreased as the electron-donating ability of the substituent on Ar² decreased (3aa-3ia). Substrates with an ortho- or *meta*-substituted Ar² ring also showed fairly good enantioselectivities (3ja, 3ka). We also investigated how substituents on Ar^1 affected the enantioselectivity when Ar^2 had an electron-donating para-methoxy group (3ma-3Aa, 3Ca). Most of the B-H bond insertion products were obtained with ee values higher than 90%. We noticed that the enantioselectivity was enhanced by increasing the electronwithdrawing ability of the substituent on Ar¹ (**3ma–3ua**). The structures and absolute configurations of 3ba and 3ra were determined by X-ray diffraction analysis of single crystals.^[41] Interestingly, when Ar¹ had an *ortho* substituent, the enantioselectivity was only slightly affected by the other substitu-



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Scheme 2. Rh-catalyzed asymmetric B–H bond Insertion of diarylcarbenes: substrate scope.^[a] [a] Reaction conditions (procedure A): Rh₂(S-TBPTTL)₄/**1/2 a** = 0.002/0.2/0.24 (mmol), PhMe, -40° C, 0.5–12 h. Isolated yields were given. The *ee* values were determined by chiral HPLC analysis. For substrate **1A**, the reaction was performed at 0°C. [b] Reaction conditions (procedure B): hydrazone (0.2 mmol), activated MnO₂ (1 mmol), MgSO₄ (0.24 mmol), DCM, 0°C, 2–3 h; then **2a** (0.24 mmol), Rh₂(S-TBPTTL)₄ (0.002 mmol), PhMe, -40° C, 6 h. [c] Reaction conditions (procedure C): hydrazone (0.2 mmol), activated MnO₂ (1 mmol), MgSO₄ (0.24 mmol), DCM, 0°C, 2–3 h; then **2a** (0.24 mmol), MgSO₄ (0.24 mmol), DCM, 0°C, 2–3 h; then **2a** (0.24 mmol), Rh₂(S-TBPTTL)₄ (0.002 mmol), PhMe, 0°C, 6 h.

ents on Ar¹ or Ar² (**3za–3Ca**). Diazo substrates containing other aryl groups, including naphthyl (**3Da**), piperonyl (**3Ea**), benzofuryl (**3Fa**), indolyl (**3Ga**), and pyridinyl (**3Ha**), also exhibited excellent enantioselectivity. Notably, all the B–H bond insertion products were stable during purification operations (e.g., chromatography and recrystallization) and could be stored for several months without any decomposition. The high stability of the products may be attributable to their coordinatively saturated boryl groups.

The synthetic potential of this B-H bond insertion reaction was then examined (Scheme 3). The reaction could be conducted at a gram scale with a catalyst loading of 0.05 mol% without compromising either the yield or the enantioselectivity, and the ee value of B-H bond insertion product (S)-3ra could be improved by crystallization (Scheme 3 A). The turnover number reached 14 300, which, to our knowledge, is the highest turnover number reported for a B-H bond insertion reaction promoted by a molecular catalvst.^[23-33,42-48] Furthermore, borane (S)-3ra could easily be converted into widely used boron reagents: chiral boronic ester 4, potassium trifluoroborate 5, and MIDA (methyliminodiacetic acid) borate 6 (Scheme 3B). Interestingly, we accidentally discovered that reaction of (S)-3ra with Nchlorosuccinimide resulted in a nearly quantitative yield of unprecedented chlorinated boron 7. In addition, (S)-3ra could be transformed to alcohol 8 by oxidation with H_2O_2 , to dibenzylic amine 9 via a potassium trifluoroborate intermediate in one pot,^[49] and to chiral triaryl methanes **10** and **11** by means of sp^2 - sp^3 coupling reactions.^[18,20] In most cases, the stereochemistry was well retained during the transformations



Scheme 3. Gram-scale experiments and transformations of product **3** ra. Reaction conditions: a) pinacol, PhMe, 90°C, 1 h; b) KHF₂ (aq.), dioxane, reflux, 2 h; c) *N*-methyl imidodiacetic acid, toluene/ DMSO = 5:1, 110°C, 17 h; d) *N*-chlorosuccinimide, THF, 1 h; e) H₂O₂, MeOH, reflux, 17 h; f) KHF₂ (aq.), dioxane, reflux, 2 h; then SiCl₄, DCE, 0°C-rt, 1 h; then BnN₃, DCE, 80°C, 1 h; then NaOH (aq.); g) 2,2-dimethyl-1,3-propanediol, PhMe, 100°C, 2 h; then, 4-iodoacetophenone, Pd(PPh₃)₄, Ag₂O, K₂CO₃, Et₂O, 60°C, 17 h; h) pinacol, PhMe, 100°C, 1 h; then 2-lithiofuran, -78°C, 1 h; then *N*-bromosuccinimide, -78°C, 1 h.

as described in the literatures,^[18,20,49] showing the potential utility of this protocol.

To elucidate the reaction mechanism, we carried out both a competitive kinetic isotope effect (KIE) experiment $(k_{\rm H}/$ $k_{\rm D} = 1.97$) and a parallel KIE experiment $(k_{\rm H}/k_{\rm D} = 5.46)$ involving the reaction between 1a and 2a (Scheme 4A and B, respectively). The large, primary KIE indicates that B-H bond cleavage might be involved in the rate-limiting step. To gain deeper insight into the mechanism, we determined the reaction order of every component by using in situ IR to measure the initial reaction rate at a series of concentrations of each component (Scheme 4C). These experiments indicated that the kinetics were first order for the Rh₂(S-TBPTTL)₄ catalyst, zero order for diazomethane 1a, and negative first order for borane adduct 2a. The kinetics profiles suggest that decomposition of 1a is not the rate-determining step and that 2a might inhibit decomposition of 1a by binding to the active site of the dirhodium catalyst.^[50] The interaction of the borane adduct with the dirhodium catalyst was confirmed by the fact that changes in the UV spectra of Rh₂(S-TBPTTL)₄ in toluene were observed upon addition of 2a (Figure S1). The pre-equilibrium formation of a restingstate complex of $Rh_2(S$ -TBPTLL)₄ and **2a** might contribute to the striking difference between the competitive and parallel $k_{\rm H}/k_{\rm D}$ values. Reactions of a mixture of borane adducts in one



Scheme 4. Control experiments.

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pot gave enantioselectivities similar to those obtained for reactions in separate pots, suggesting that only one molecule of the borane adduct is involved in the enantio-determining step (Scheme 4D).

Furthermore, we conducted density functional theory calculations (Gaussian 09) on the rhodium-catalyzed reaction of diazomethane 1r and amine-borane adduct 2a (Scheme 5A). The calculations indicate that the catalytic cycle starts with dissociation of one borane adduct from [Rh·2a·2a] (a process that is uphill by $6.6 \text{ kcal mol}^{-1}$). Diazomethane **1r** coordinates to the rhodium catalyst by replacing 2a and then decomposes to release N2 and carbene intermediate CB via transition state TS1, a process with an activation energy of only 11.5 kcal mol⁻¹. Once formed, **CB** can insert into the B-H bond of 2a via three-membered-ring transition state TS2-S, generating the desired product and releasing the catalyst to start another catalytic cycle. The B-H bond insertion process has an activation energy of 16.3 kcal mol⁻¹, which is 4.8 kcal mol⁻¹ higher than the energy for diazo decomposition. These results indicate that B-H insertion is the rate-determining step, which agrees well with the results of the KIE experiments and with the zero order kinetics observed for diazo-



Scheme 5. Calculated catalytic cycle,^[a] chiral control model,^[b] and Hammett plots.^[c] [a] Calculated catalytic cycle of rhodium-catalyzed B⁻⁻ H insertion of **1** r and **2** a. Density functional theory calculations were performed at the B3LYP-D3(BJ)/DEF2TZVP/SMD(PhMe)//B3LYP/ DEF2SVP. Gibbs free energies relative to **[Rh]** were given in the bright blue box. [b] Optimized lowest-energy transition structures for *R* and *S* products. Gibbs free energies were given relative to **TS2-S**. [c] Hammett studies. Hammett parameters σ and σ^+ were used for electron-withdrawing group (EWG-) and electron-donating group (EDG-) substituted arenes, respectively. Other correlation trials were given in Tables S3–S5 in the Supporting Information.

methane 1a (Scheme 4A-C). The irreversible B-H bond insertion can be considered as the enantio-determining step. The structures corresponding to the lowest-energy transition states for the major and minor enantiomers of the product are presented in Scheme 5 B. In these two transition states TS2-R and TS2-S, the p-OMe aryl ring tends to adopt a more coplanar orientation with carbene p orbital, while p-Cl aryl ring adopts a tilted conformation. Such orientation difference of two aryl rings causes steric difference in asymmetrical environment of catalyst.^[38-40] In accord with experimental observations, the calculated energy of TS2-S was 2.1 kcal mol⁻¹ lower than that of **TS2-R**, which leads to the disfavored enantiomer, (R)-3ra. This energy difference might arise from steric repulsion between the p-OMe-substituted arene and the carboxylate group of the rhodium catalyst in TS2-R, repulsion that is absent in TS2-S.

Hammett analysis established that the enantioselectivity, ln(er), was linearly related to the electronic difference $(\sigma_1 - \sigma_2^+)$ between the two arene rings of the diazo substrates (Scheme 5 C). This result is consistent with the buildup of positive charge on the carbene carbon after decomposition of the diazomethane. The positive charge buildup causes the different behavior of the two aryl rings: the electron-deficient ring tends to be tilted out of the plane of the rhodium carbene, whereas the electron-rich ring tends to be coplanar with the carbene.^[38,40,51] The greater the electronic difference between the two arenes is, the better the enantioselectivity will be.

Conclusion

We have realized a method for rhodium-catalyzed chiral B-H bond insertion reactions of borane adducts and diaryl diazomethanes. Using this method, we prepared various gemdiarylmethine borane compounds in high yield with excellent enantioselectivity, and the mild reaction conditions showed good functional group tolerance. The B-H insertion products could easily be transformed to widely used pinacolborates, potassium trifluoroborates, MIDA borates, diarvl methanol compounds, diaryl methyl amines, and triaryl methanes. Kinetics experiments and density functional theory calculations suggest that the borane adduct coordinates to the rhodium catalyst and interferes with decomposition of the diazomethane and that insertion of the rhodium carbene into the B-H bond is the rate-determining step (rather than diazomethane decomposition). Hammett plots showed that the enantioselectivity of the reaction correlates to electronic differences between the two aryl groups of the diazomethane. This method not only opens up a new route to gem-diarylmethine boranes but also deepens our understanding of B-H bond insertion reactions.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords: B–H bond insertion \cdot chiral dirhodium catalysts \cdot diaryl carbenes \cdot diaryl diazomethanes \cdot gem-diarylmethine boranes

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